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SHEET 1 OF 3

SUBSTITU	TE FO	R FORM IPO TO THE STATE OF THE	ATTY DOCKET NO: FOGH=5A		SERIAL NO: 10/588,082			
INFORMATION DISCLOSURE STATEMENT LIST OF DOCUMENTS CITED BY APPLICANT			FIRST INVENTOR: FOGH, Jens					
			FILING DATE: June 20, 2007					
			EXAMINER:		ART UNIT:			
U.S. PATENT DOCUMENTS (include at least patentee, patent/pub number and filing/issue/pub date)								
EXAM. INITIAL	ID	DOCUMENT NUMBER	FILING, ISSUE OR PUBLICATION DATE MM-DD-YYYY	PATENTEE OR APPLICANT	Relevant Passage(s)	Ť		
FOREIGN I	PATEN	IT DOCUMENTS (include a	nt least document nur	mber, publication date and c	ountry)			
EXAM. INITIAL	ID	COUNTRY CODE & DOCUMENT NUMBER	PUBLICATION DATE MM-DD-YYYY	PATENTEE OR APPLICANT	Relevant Passage(s)	т*		
OTHER DOCUMENTS (include AUTHOR, title, name of publication, volume, pages & date of publication) Please list in alphabetical order.								
	AK	Baum, et al., "The assay of arylsulphatases A and B in human urine", <u>Clin. Chim. Acta,</u> vol. 4, pp 453-455, 1959.						
	AL	Coenen, et al. "Morphological alterations in the inner ear of the arylsulfatase A-deficient mouse", <u>Acta Neuropathol</u> , Vol. 101, pp 491-498, 2001.						
	ΑM	Demeule, et al. "High transcytosis of melanotransferrin (P97) across the blood-brain barrier", <u>Journal of Neurochemistry</u> , Vol. 83, pp 924-933, 2002.						
	AN	D'Hooge et al., "Hyperactivity, neuromotor defects, and impaired learning and memory in a mouse model for metachromatic leukodystrophy", <u>Brain Research</u> , Vol. 907 pp 35-43, 2001.						
	AO	Dierks et al., "Conversion of cysteine to formylglycine: A protein modification in the endoplasmic reticulum", Proc. Natl. Acad. Sci. USA, Vol. 94, pp 11963-11968, October 1997.						
	AP	Dunican, et al., "Designing Cell-Permeant Phosphopeptides to Modulate Intracellular Signaling Pathways", Biopolymers (Peptide Science), Vol. 60 pp 45-60, 2001.						
	AQ	Gieselmann, et al., "Metachromatic leukodystrophy: Molecular genetics and an animal model", <u>J.</u> Inher. Metab. Dis., Vol. 21, pp 564-574, 1998.						
	AR		mann, et al., "Metachromatic leukodystrophy: consequences of sulphatide accumulation, Acta atr Suppl., Vol. 443, pp 74-79, 2003.					
EXAMINER				DATE CONSIDERED				
<b>EXAMINER:</b> Initial if reference considered. Draw line through citation if not in conformance <u>and</u> not considered. Include copy of this form with next communication to applicant.								

<sup>\* &</sup>quot;Relevant Passages" column is optional. Put check in "T" column if English translation of entire document included. If English language abstract included, put A in this box. If ref. in English, put "E". If requirement otherwise met, put O.

SHEET 2 OF 3

SUBSTITUTE FOR FORM IPC/SB/08 U.S. DEPARTMENT OF COMMERCE PATENT AND TRADEMARK OFFICE			ATTY DOCKET NO: FOGH=5A	SERIAL NO: 10/588,082		
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	AS	V. Gieselmann, et al., "In Vitro Mutagenesis of Potentioal N-Glycosylation Sites of Arylsulfatase A", Journal of Biological Chemistry, Vol. 267, No. 19, pp 13262-13266, July 5, 1992.				
	АТ	Hess, et al., "Phenotype of arylsufatase A-deficient mice: Relationship to human metachromatic leukodystrophy", Proc. Natl. Acad. Sci. USA, Vol. 93, pp 14821-14826, December 1996.				
	AU	Ho, et al., "Synthetic Protein Transduction Domains: Enhanced Transduction Potential in Vitro and in Vivo <sup>1</sup> ", Cancer Research, Vol. 61, pp 474-477, January 15, 2001.				
	AV	Kudoh, et al., "Diagnosis of Metachromatic Leukodystrophy, Krabbe Disease, and Farber Disease after Uptake of Fatty Acid-labeled Cerebroside Sulfate into Cultured Skin Fibroblasts", <u>J. Clin. Invest.,</u> Vol. 70, pp 89-97, July 1982.				
	AW	Lindgren, et al., "Cell-penetrating peptides", <u>TIPS</u> , Vol. 21, pp. 99-103, March 2000.				
	AX	Lukatela, et al., "Crystal Structure of Human Arylsufatase A: The Aldehyde Function and the Metal Ion at the Active Site Suggest a Novel Mechanism for Sulfate Ester Hydrolysis", Biochemistry, Vol. 37, pp 3654-3664, 1998.				
	AY	Lüllmann-Rauch, et al., "Lysosomal sulfoglycolipid storage in the kidneys of mice deficient for arysulfatase A (ASA) and of double-knockout mice deficient for ASA and galactosylceramide synthase", <u>Histochem Cell Biol</u> , Vol. 116, pp 161-169, 2001.				
	AZ	Matsushima, et al. "Absence of MHC Class II Molecules Reduces CNS Demyelination, Microglial/Macrophage Infiltration, and Twitching in Murine Globoid Cell Leukodystrophy", <u>Cell</u> , Vol. 78, pp 645-656, August 26, 1994.				
	ВА	Matzner, et al., "Long-term expression and transfer of arylsulfatase A into brain of arylsulfatase A-deficient mice transplanted with bone marrow expressing the arylsulfatase A cDNA from a retroviral vector", Gene Therapy, Vol. 7, pp 1250-1257, 2000.				
	вв	Matzner et al., "Retrovirally expressed human arylsufatase A corrects the metabolic defect of arylsulfatase A-deficient mouse cells", Gene Therapy, Vol. 7, pp 805-812, 2000.				
	вс	Muschol, et al., "Secretion of phosphomannosyl-deficient arylsulphatase A and cathepsin D from isolated human macrophages", Biochem J., Vol. 368, pp 845-853, 2002.				
	BD	Pan et al., "TNF <i>a</i> Transport across the Blood-Brain Barrier is Abolished in Receptor Knockout Mice", Experimental Neurology, Vol. 174, pp 193-200, 2002.				
•	BE	Pan et al., "Upregulation of the Transport System for TNF <i>α</i> at the Blood-Brain Barrier", <u>Archives of Physiology and Biochemistry</u> , Vol. 109, No. 4, pp 350-353, 2001.				
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	BF	Rodman, et al., Circulating natural IgM antibodies and their corresponding human cord blood cell-derived Mabs specifically combat the Tat protein of HIV", Experimental Hematology, Vol. 29 pp 1004-1009, 2001.				
	ВG	Rothenberger, et al., "Coincident expression and distribution of melanotransferrin and transferring receptor in human brain capillary endothelium", <u>Brain Research</u> , Vol. 712, pp 117-121, 1996.				
	вн	Sandhoff, et al., "Kidney Sulfatides in Mouse Models of Inherited Glycosphingolipid Disorders", <u>The Journal of Biological Chemistry</u> , Vol. 277, No. 23, pp 20386-20398, 2002.				
	ві	Schmidt, et al., "A Novel Amino Acid Modification in Sulfatases That is Defective in Multiple Sulfatase Deficiency", Cell, Vol. 82, pp 271-278, July 28, 1995.				
	BJ	Schwarze, et al., Protein transduction: unrestricted delivery into all cells?", trends in CELL BIOLOGY, Vol. 10, pp 290-295, July 2000.				
	вк	Scott, et al., "Differential Staining of Acid Glycosaminoglycans (Mucopolysaccharides) by Alcian Blue in Salt Solutions", <u>Histochemie</u> , Vol. 5, pp 221-233, 1965.				
	BL	Selmer, et al., "The evolutionary conversation of a novel protein modification, the conversion of cysteine to serinesemialdehyde in arylsufatase from Volvox carteri", Eur. J. Biochem., Vol. 238, pp 341-345, 1996.				
_	вм	Sommerlade, et al., "Four monoclonal antibodies inhibit the recognition of arylsulphatase A by the lysosomal enzyme phosphotransferase", Biochem J., Vol. 297, pp 123-130, 1994.				
	BN	Wada, et al., "Microglial activation precedes acute neurodegeneration in Sandhoff disease and is suppressed by bone marrow transplantation", <a href="Proc. Nat. Acad. Sci. (USA)">Proc. Nat. Acad. Sci. (USA)</a> , Vol. 97, No. 20, pp 10954-10959, September 26, 2000.				
	во	Wittke, et al., "Lysosomal sulfatide storage in the brain of arylsulfatase A-deficient mice: cellular alterations and topographic distribution", Acta Neurophatol, Vol. 108, pp 261-271, 2004.				
	ВР	Wu, et al., "Neuroprotection with noninvasive neurotrophin delivery to the brain", <u>Proc. Natl. Acad. Sci.</u> USA, Vol. 96, pp 254-259, January 1999.				
	BQ	Yao, et al., "Microanalysis of Complex Tissue Lipids by High-Performance Thin-Layer Chromatography", Analytical Biochemistry, Vol. 150, pp 111-116, 1985.				
	BR	Zielasek, et al., "Functional Abnormalities in $P_0$ -Deficient Mice Resemble Human Hereditary Neuropathies Linked to $P_0$ Gene Mutations", Muscle & Nerve, Vol. 19, pp 946-952, 1996.				
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/Yong Pak/

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